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(54) Title: RADIATION-CURABLE COMPOSITIONS	HAVIN	IG FAST CURE SPEED AND GOOD ADHESION TO GLASS

(57) Abstract

A radiation-curable composition for use as an optical fiber material or coating comprising the combination of the following pre-mixture ingredients: (A) about 5 wt.% to about 95 wt.% of one or more radiation-curable oligomers; (B) about 5 wt.% to about 95 wt.% of one or more monomer diluents, (C) optionally, one or more photoinitiators, (D) about 0.1 wt.% to about 30 wt.% of one or more about promoters represented by: R¹-L-Si(OR); wherein R¹ is a group comprising an ethylenically unsaturated radiation-curable group; and said three R² groups independently of each other are C₁-C₁₀ groups; and L is a linking group which comprises one or more alknoy or branched propoxy groups.

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- 1 -

5 RADIATION-CURABLE COMPOSITIONS HAVING FAST CURE SPEED AND GOOD ADHESION TO GLASS

FIELD OF THE INVENTION

This invention relates to radiation-curable

compositions, and in particular, to compositions which

are useful as optical fiber coatings and have both good

adhesion to glass and fast cure speed.

DESCRIPTION OF RELATED ART

Radiation-curable compositions are extensively used by the optical fiber industry for the production of optical fibers, ribbons, and cables. For example, optical glass fibers are routinely coated with a protective radiation-curable coating system

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- 20 immediately after the glass fiber is manufactured in a draw tower furnace so as to preserve the pristine character of the freshly manufactured glass fiber. Immediately after a radiation-curable coating is applied to protect the fiber, the coating is rapidly
- 25 cured by exposure to radiation (commonly ultraviolet light). The industry demands faster production speeds, and therefore, faster radiation-curing coating compositions.
- Usually, two radiation-curable coatings are
 needed to maximize fiber performance. A soft inner
 primary coating directly contacts the glass fiber and
 helps prevent microbending, and a harder outer primary
 coating overlays the soft inner primary coating and
 protects it during handling. Single coating systems are
 twown but are not generally used at the present.

Radiation-curable matrix and bundling materials can further support and protect the individual strands of coated fiber when individual strands are bundled together into optical fiber 5 ribbons, cables, and associated structures. Also, radiation-curable inks can be used to color code individual strands of optical fiber. All of these types of optical fiber-related materials are preferably radiation-curable, and fast cure for each is essential.

The adhesion of different optical fiber materials to one another is also very important. However, combinations of properties can be difficult to achieve.

Examples of radiation-curable inner primary
15 coatings are disclosed in U.S. Patent No. 5,336,563 to
Coady et al and of outer primary coatings in U.S.
Patent No. 4,472,019 to Bishop et al. Additional
aspects of optical fiber coating technology are
disclosed in, for example, U.S. Patent Nos. 5,595,820
20 to Szum; 5,199,098 to Nolan et al.; 4,923,915 to Urruti
et al.; 4,720,529 to Kimura et al.; and 4,474,830 to
Taylor et al.

To maximize cure speed in an ultraviolet light cure, at least one photoinitiator is required (although photoinitiator may be omitted in an electron beam cure). Several photoinitiators can be used to achieve a suitable balance of surface and through cure. Mono-acyl phosphine oxide type photoinitiators can be used such as Lucirin TPO (2,4,630 trimethylbenzoyl)diphenyl phosphine oxide, which exhibits relatively fast cure speed but can also generate undesirable crystallization effects. Other phosphine oxide photoinitiators can show reduced amounts of harmful crystallization effect, but they may

35 also have slower cure speed or cause yellowing. This illustrates again how it can be difficult to achieve

35

fast cure speed in combination with other desirable properties.

Fast cure is particularly difficult to achieve with inner primary coatings. For these coatings, fast-cure must not be achieved without loss of adhesion to the glass fiber. U.S. Patent No. 4,849,462 to Bishop teaches that the silane adhesion promoter, mercaptopropyltrimethoxy silane, is particularly useful for improving adhesion to optical glass fiber. Adhesion promotion is particularly important under wet aging conditions. However, despite their advantages, the mercaptosilane and other typical silane adhesion promoters also tend to decrease cure speed. The mercaptosilane adhesion promoter also tends to reduce coating modulus and liquid coating shelf life.

By way of example, the aforementioned U.S. Patent No. 5,336,563 to Coady et al discloses use of silane adhesion promoters and cure speed measurements for inner primary coatings. According to this patent, cure speeds of 0.45, 0.32, and 0.37 J/cm (measured as dose at 95% maximum attainable modulus) are rapid based on industrial standards. This patent, however, does not teach or suggest how to prepare compositions having cure speed of less than about 0.2 J/cm. More specifically, this patent does not teach or suggest how to attain both fast cure speed and good wet adhesion.

SUMMARY OF THE INVENTION

An object of the present invention is to provide radiation-curable compositions having both fast cure speed and good adhesion to glass. An additional object is to provide compositions having both fast cure speed and good interlayer adhesion.

These and other objects are achieved by a

radiation-curable composition for use as an optical fiber coating or material comprising the combination of the following pre-mixture ingredients:

(A) about 5 wt.% to about 95 wt.% of one or

- 5 more radiation-curable oligomers,
 - (B) about 5 wt.% to about 95 wt.% of one or more monomer diluents,
 - (C) optionally, one or more photoinitiators.
- (D) about 0.1 wt.* to about 30 wt.* of one or more adhesion promoters represented by: R*-L-Si(OR*).

wherein R¹ is a group comprising an ethylenically unsaturated radiation-curable group; wherein said three 5 R² groups independently of each other are C₁-C₁₀ groups; and wherein L is a linking group which comprises one or more branched propoxy groups.

The present invention also provides radiation-cured forms of the radiation-curable composition, as well as coated optical fibers comprising this radiation-cured form. The present invention also provides optical fiber cable and ribbon assemblies and methods of making coated optical fiber.

25 DETAILED DESCRIPTION OF THE PRESENT INVENTION

Known examples of radiation-curable compositions for optical fiber materials and coatings can be found in, for example, U.S. Patent Nos. 5,384,342; 5,456,984; 5,596,669; 5,336,563; 5,093,386; 4,716,209; 4,624,994; 30 4,572,610; and 4,472,019. These references teach one skilled in the art how to formulate and use compositions based on acrylate-functionalized oligomers and monomer diluents.

For this invention, "pre-mixture

35 ingredient" means an ingredient before it is mixed with

other ingredients. When formulating a radiation-curable composition from its pre-mixture ingredients, some interaction or reaction of the ingredients may be possible after mixing. However, pre-mixture ingredient refers to the identity of the ingredient before mixing. The present invention is not generally limited by the order in which the pre-mixture ingredients are mixed.

" (Meth) acrylate" means acrylate.

methacrylate, or a mixture thereof.

10 The oligomer system A comprises one or more radiation-curable oligomers. The radiation-curable oligomer(s) A can be any known radiation-curable oligomer used in radiation-curable optical fiber material or coating compositions. An example of a 15 suitable radiation-curable oligomer A includes a urethane oligomer having a molecular weight of at least about 500 and containing at least one ethylenically unsaturated group that can be polymerized through actinic radiation. Preferably, the oligomer A is 20 substantially free of isocyanate functionality. Preferably, the oligomer A has two terminal radiationcurable functional groups, one at each end of the oligomer. Urethane-free oligomers may also be used in this composition.

25 Preferably, the number average molecular weight of the oligomer A is higher than about 700 or more, preferably the molecular weight is higher than about 1,000 and most preferably higher than about 2,000. Preferably when acrylated acrylics are used, the 30 number average moleculaar weight is lower than about 50,000, and for other oligomers lower than about 10,000.

The radiation-curable oligomer system A is preferably present in an amount of about 5 to about 95 35 % by weight, and more preferably, about 20 to about 80% by weight. If multiple oligomers are used, the amounts

of each oligomer are added to determine the amount of oligomer system A. All weight percentages used herein are relative to the total weight of the composition unless indicated otherwise.

Examples of suitable radiation-curable functional groups for oligomer(s) A include ethylenically unsaturated groups having (meth)acrylate. vinylether, acrylamide, maleimides, maleate or fumarate functionality. Preferably, the radiation-curable group 10 in the oligomer A is a (meth)acrylate or vinylether group. Most preferably, the radiation-curable group is

Preferably, the oligomer A contains at least two ethylenically unsaturated groups which are 15 bound to an oligomer backbone. For example, ethylenically unsaturated groups can be present at each end of the oligomer backbone as reactive termini.

The oligomer backbone can comprise one or more polymer blocks of repeating units coupled with 20 each other via, for example, urethane linkages. Methods to allow for such coupling are known in the art. For example, oligomer A can be prepared by reaction of (i) an oligomer polyol, (ii) a diisocyanate compound, and (iii) a hydroxy functional ethylenically unsaturated monomer such as, for example,

hydroxyalkyl (meth) acrylate.

an acrylate group.

Preferably, the oligomer backbone comprises repeat units of polyether, polyolefin, polyester, polycarbonate, or mixtures thereof. Alternatively, 30 acrylated acrylic type oligomers can be used. If the oligomer backbone is a polyether, the resulting materials generally have a low glass transition temperature and good mechanical properties. If the oligomer backbone is a polyolefin, the resulting 35 materials generally have further improved water resistance.

- 7 -

If an oligomer backbone polyol is used, preferably it has on average at least about 2 hydroxyl groups. The oligomer backbone polyol may have, on average, more than 2 hydroxyl groups. Examples of such 5 an oligomer diol include polyether diols, polyolefin diols, polyester diols, polycarbonate diols, and mixtures thereof. Polyether and polyolefin diols, or combinations thereof, are preferred.

If a polyether diol is used, preferably the 0 polyether is a substantially non-crystalline polyether. Preferably, the polyether comprises repeating units of one or more of the following monomer groups: -0-CH3-CH3-

-O-CH2-CH (CH3) -

15 -O-CH2-CH2-CH2-

-O-CH2-CH2-CH2-CH2-

-O-CH2-CH (CH3) -CH2-CH2-

-O-CH(CH₃)-CH₂-CH₂-CH₂--OCH₂C(CH₃)₂-

20 -O-CH (CH2-CH3) -CH2-.

Hence, the polyether can be made from epoxy-ethane, epoxy-propane, tetrahydrofuran, methylsubstituted tetrahydrofuran, epoxybutane, and the like.

25 An example of a polyether polyol that can be used is the polymerization product of 20 percent by weight of 3-methyltetrahydrofuran and 80 percent by weight of tetrahydrofuran, both of which have undergone a ring opening polymerization. This polyether copolymer contains both branched and non-branched oxyalkylene repeating units and is marketed as PTG-L 1000 (Hodogaya Chemical Company of Japan). Another example of a polyether that can be used is PTG-L 2000 (Hodogaya Chemical Company).

35 If a polyolefin diol is used, the polyolefin is preferably a linear or branched

hydrocarbon containing a plurality of hydroxyl end groups. Preferably, the hydrocarbon is a non-aromatic compound containing a majority of methylene groups (-CH2-) and which can contain internal unsaturation and/or 5 pendent unsaturation. Fully saturated, for example. hydrogenated hydrocarbons, are preferred because the long term stability of the cured optical fiber coating increases as the degree of unsaturation decreases. Examples of hydrocarbon diols include, for example, 10 hydroxyl-terminated, fully or partially hydrogenated 1,2-polybutadiene: 1,4-1,2-polybutadiene copolymers, 1,2-polybutadiene-ethylene or -propylene copolymers, polyisobutylene polyol; mixtures thereof, and the like. Preferably, the hydrocarbon diol is a substantially, 15 fully hydrogenated 1,2-polybutadiene or 1,2polybutadiene-ethene copolymer.

Examples of polycarbonate diols are those conventionally produced by the alcoholysis of diethylene carbonate with a diol. The diol can be, for 20 example, an alkylene diol having about 2 to about 12 carbon atoms, such as, 1,4-butane diol, 1,6-hexane diol, 1,12-dodecane diol, and the like. Mixtures of these diols can also be utilized. The polycarbonate diol can contain ether linkages in the backbone in 25 addition to carbonate groups. Thus, for example, polycarbonate copolymers of alkylene oxide monomers and the previously described alkylene diols can be used. Alkylene oxide monomers include, for example, ethylene oxide, tetrahydrofuran, and the like. These copolymers 30 produce cured coatings that exhibit a lower modulus and also inhibit crystallinity of the liquid coating composition compared to polycarbonate diol homopolymers. Admixtures of the polycarbonate diols and polycarbonate copolymers can also be utilized. 35

Polycarbonate diols include, for example,
Duracarb 122 (PPG Industries) and Permanol KM10-1733

(Permuthane, Inc., Ma.). Duracarb 122 is produced by the alcoholysis of diethylcarbonate with hexane diol. Examples of polyester diols include the

reaction products of saturated polycarboxylic acids, or 5 their anhydrides, and diols. Saturated polycarboxylic acids and anhydrides include, for example, phthalic acid, isophthalic acid, terephthalic acid, trimellitic acid, tetrahydrophthalic acid, hexahydrophthalic acid, tetrachlorophthalic acid, adipic acid, azelaic acid,

sebacic acid, succinic acid, glutaric acid, malonic acid, pimelic acid, suberic acid, 2,2-dimethylsuccinic acid, 3,3-dimethylglutaric acid, 2,2-dimethylglutaric acid, the like, anhydrides thereof and mixtures thereof. Diols include, for example, 1,4-butanediol,

15 1,8-octane diol, diethylene glycol, 1,6-hexane diol, dimethylol cyclohexane, and the like. Included in this classification are the polycaprolactones, commercially available from Union Carbide under the trade designation Tone Polylol series of products, for

20 example, Tone 0200, 0221, 0301, 0310, 2201, and 2221.
Tone Polyol 0301 and 0310 are trifunctional.

One or more polyisocyanates can also be used to prepare oligomer A. Any known organic polyisocyanate (ii), alone or in admixture, can be used 25 as the polyisocyanate. However, aliphatic, non-aromatic polyisocyanates are preferred if non-yellowing is to be achieved

Generally, in oligomer synthesis, an oligomer product is obtained which is end-capped with 30 the reaction product from the isocyanate/ethylenically unsaturated monomer reaction on at least one end of the oligomer molecule. "End-capped" means that a functional group caps an end of the oligomer diol.

The isocyanate/hydroxy functional monomer

35 reaction product attaches to the oligomer backbone (i)
diol via a urethane linkage. The urethane reactions can

take place in the presence of a catalyst. Catalysts for the urethane reaction include, for example, diazabicyclooctane crystals and the like.

Preferably the polyisocyanate (ii) is a 5 diisocyanate. Examples of diisocyanates (ii) include tetramethylxylylene diisocyanate (TMXDI), isophorone diisocyanate (IPDI), toluene diisocyanate (TDI), diphenylmethylene diisocvanate, hexamethylene diisocyanate, cyclohexylene diisocyanate, methylene

10 dicyclohexane diisocyanate, 2,2,4-trimethyl hexamethylene diisocyanate, m-phenylene diisocyanate, 4-chloro-1,3-phenylene diisocyanate, 4,4'-biphenylene diisocyanate, 1,5-naphthylene diisocyanate, 1,4tetramethylene diisocvanate. 1.6-hexamethylene

15 diisocyanate, 1,10-decamethylene diisocyanate, 1,4cyclohexylene diisocyanate, and polyalkyloxide and polyester glycol diisocvanates such as polytetramethylene ether glycol terminated with TDI and polyethylene adipate terminated with TDI, respectively. 20 Preferably, the isocvanates are TDI or IPDI.

Generally the compound providing a reactive terminus (iii) contains a functional group which can polymerize under the influence of actinic radiation. and the compound contains a functional group which can 25 react with the diisocyanate. Hydroxy functional ethylenically unsaturated monomers are preferred. More preferably, the hydroxy functional ethylenically unsaturated monomer contains acrylate, (meth)acrylate, vinyl ether, maleate or fumarate functionality.

In the reaction between hydroxy group of (i) and isocyanate groups of (ii), it is preferred to employ a stoichiometric balance between hydroxy and isocyanate functionality and to maintain the reaction temperature of at least 25°C. The hydroxy functionality 35 should be substantially consumed. The hydroxy functional ethylenically unsaturated monomer attaches

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to the isocyanate via a urethane linkage. Monomers having (meth)acrylate functional groups include, for example, hydroxy functional acrylates such as 2-hydroxyethyl acrylate, 2-hydroxypropyl acrylate, and 5 the like. Monomers having vinyl ether functional groups include, for example, 4-hydroxybutyl vinyl ether, and triethylene glycol monovinyl ether. Monomers having maleate functional groups include, for example, maleic acid and hydroxy functional maleates.

Alternatively, the radiation-curable oligomer A can be an acrylic oligomer comprising at least one radiation-curable (meth)acrylate group, and preferably, at least one acrylate group. These are known in the art as acrylated acrylics.

The invention is not believed to be limited by how the acrylated acrylic oligomer, or any other oligomer, is prepared. Oligomer synthetic routes for acrylated acrylics can, for example, involve an esterification of a hydroxyl-functional acrylic oligomer with

20 (meth)acrylic acid, or the reaction of an epoxyfunctional acrylic oligomer with (meth)acrylic acid. These acrylated acrylics can include urethane linkages.

Acrylic monomers which can be used to prepare the acrylic oligomer can be represented as the 25 esters represented in formula (1).

 $CH_2 = CHCOOR$ (1)

In formula (1), the acrylic monomers can be various types of esters including, for example, n-alkyl esters, secondary and branched-chain alkyl esters, esters of olefinic alcohols, aminoalkyl esters, esters of ether alcohols, cycloalkyl esters, and esters of halogenated alcohols, glycol diacrylates.

Methacrylic monomers, $CH_2 = C(CH_3)COOR$, which are analogous to those of the acrylic monomers in formula (1) can also be used. In general, monomers represented as $CH_2 = CR_2COOR$ can be used wherein R_1 is a C_1 - C_6 alkyl.

N-alkyl esters in formula (1) include R being methyl, ethyl, propyl, butyl, pentyl, hexyl, heptyl, octyl, nonyl, decyl, dodecyl, tetradecyl, hexadecyl:

secondary and branched-chain alkyl esters in formula (1) include R = isopropyl, isobutyl, secbutyl, 2-ethylhexyl, 2-ethyl butyl;

esters of olefinic alcohols in formula (1)
include R = allyl, 2-methylallyl, furfuryl, 2-butenyl;
10 aminoalkyl esters in formula (1) include R
= 2-(dimethylamino)ethyl, 2-(diebutylamino)ethyl, 2(dibutylamino)ethyl, and 3-(diethylamino)propyl;

esters of ether alcohols include R = 2-methoxyethyl, 2-ethoxyethyl, tetrahydrofurfuryl, 2-

methoxyethyl; 2-ethoxyethyl, tetranydrolulldryl, 215 butoxyethyl;

cycloalkyl esters include R = cyclohexyl,
4-methylcyclohexyl, 3,3,5-trimethylcyclohexyl, 3,3,5trimethylcyclohexyl, styrenes and vinyl acetate;
esters of haloemated alcohols include R =

25 dipropylene glycol, 2,5-hexanediol, 2,2-diethyl-1,3propanediol, 2-ethyl-1,3-hexanediol, and 1,10decanediol

Additional (meth)acrylic acid and ester polymers are disclosed in, for example, Encyclopedia of

30 Polymer Science & Engineering, Vol. 1, 1985, pgs. 211305, the complete disclosure of which is hereby incorporated by reference.

Acrylated acrylics can be prepared by known synthetic methods including, for example, (1) partial 35 esterification of acrylic polymers having pendant carboxylic acid group with hydroxyethyl acrylate or

glycidyl methacrylate, or in the alternative, acrylation of glycidyl methacrylate terpolymer with acrylic acid, or (2) polymerization of monomers which already have acrylate groups such as, for example, allyl methacrylate.

The acrylic oligomer typically will have a copolymeric backbone. The Tg of the oligomer can be lowered by decreasing the content of methyl methacrylate.

The compositions according to the invention also comprise a monomer, or reactive, diluent system which comprise at least one, and preferably at least two, monomer diluent compounds. The reactive diluent can be used to adjust the viscosity of the coating composition. Thus, the reactive diluent can be a low

composition. Thus, the reactive diluent can be a low viscosity monomer containing at least one functional group capable of polymerization when exposed to actinic radiation.

The reactive diluent is preferably added in such an amount that the viscosity of the coating composition is in the range of about 1,000 to about 10,000 mPa.s. However, there is no particular limitation on viscosity, and it can be adjusted to a given application by known methods. For example,

25 viscosity can be adjusted depending on the type of optical fiber material being formulated and the method of application.

Suitable amounts of the reactive diluent system have been found to be about 5 wt% to about 95 wt%, and more preferably, about 20 wt.% to about 80 wt.%, and more preferably, about 30 wt.% to about 70 wt.%. If more than one reactive diluent is present, the amounts of reactive diluent are added together to determine the amount of the diluent system.

35 A monomer diluent preferably has a molecular weight of not more than about 550 or a viscosity at room temperature of not more than about 500 mPa.s (measured as 100% diluent).

The radiation-curable functional group present on the reactive diluent may be of the same

nature as that used in the radiation-curable oligomer.
Preferably, the radiation-curable functional group present in the reactive diluent is capable of copolymerizing with the radiation-curable functional group present on the radiation-curable oligomer.

10 Ethylenic unsaturation is preferred. In particular,

acrylate unsaturation is preferred.

Preferably, the reactive diluent system comprises a monomer or monomers having an acrylate or

vinyl ether functionality and a C₄-C₇₀ alkyl or
15 polyether moiety. Examples of such reactive diluents
include hexylacrylate, 2-ethylhexylacrylate,
isobornylacrylate, decylacrylate, laurylacrylate,
stearylacrylate, ethoxyethoxy-ethylacrylate,
laurylvinylether, 2-ethylhexylvinyl ether, N-vinyl

20 formamide, isodecyl acrylate, isooctyl acrylate, vinylcaprolactam, N-vinylpyrrolidone, acrylamide, nonylphenol acrylate and the like.

Another preferred type of reactive diluent is a compound comprising an aromatic group. Examples of 25 diluents having an aromatic group include:

ethyleneglycolphenyletheracrylate,
polyethyleneglycolphenyletheracrylate,
polypropyleneglycolphenyletheracrylate, and
alkyl-substituted phenyl derivatives of the above

30 monomers, such as

polyethyleneglycolnonylphenyletheracrylate.

Furthermore, a reactive diluent can contain two groups capable of polymerization using actinic radiation. A diluent having three or more of such such monomers include:

C₁-C₁₈ hydrocarbondioldiacrylates,
C₄-C₁₈ hydrocarbondivinylethers,
C₁-C₁₈ hydrocarbontrioltriacrylates,
the polyether analogs thereof, and
the like, such as
1,6-hexanedioldiacrylate,
trimethylolpropanetriacrylate,
hexanedioldivinylether,
triethyleneglycoldiacrylate,
pentaeritritoltriacrylate,
tripropyleneglycol diacrylate
alkoxylated bisphenol A diacrylate.

Preferably, the oligomer and the reactive diluent each contain an acrylate group as a radiation-curable group.

The composition may optionally further comprise at least one photoinitiator. Photoinitiator is required for a fast UV cure but may be omitted for electron beam cure. Conventional photoinitiators can be used. Examples include benzophenones, acetophenone derivatives, such as alpha-hydroxyalkylphenylketones, benzoin alkyl ethers and benzil ketals, monoacylphosphine oxides, and bisacylphosphine oxides. A preferred photoinitiator is 1-

25 hydroxycyclohexylphenylketone (IRGACURE 184, Ciba Geigy). Another preferred example is Lucirin TPO (BASF). Often, mixtures of photoinitiators provide a suitable balance of properties. Maleimide can be used as a photoinitiator with compositions comprising

30 acrylates. Maleimide and vinyl ethers can be photopolymerized without the need of photoinitiators.

The amount of photoinitiator system is not particularly limited but will be effective to provide fast cure speed, reasonable cost, good surface and

35 through cure, and lack of yellowing upon aging. Typical amounts can be, for example, about 0.3 wt.* to about 10

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wt.%, and preferably, about 1 wt.% to about 5 wt.%. The compositions according to the present invention further comprise one or more adhesion

promoters represented by the formula:

R1-L-Si(OR2),

wherein R1 is a group comprising one or more ethylenically unsaturated radiation-curable group; and said three R2 groups independently of each other are C1-C10 groups; and L is a linking group which comprises one 10 or more branched alkoxy groups in general and branched propoxy groups in particular. In addition, L may contain additional groups; for example, a propyl group or an aminopropylene group.

The ethylenically unsaturated radiationcurable group in R1 can be any group discussed above with respect to the radiation-polymerizable oligomers or diluents. An (meth) acrylate group is preferred, and an acrylate group is more preferred

- One or more of the R2 groups should allow 20 the propoxylated silane compound to be hydrolyzable. In particular, they should allow at least one, and preferably two or three, of the -Si-O-R2 linkages to readily react with water at room temperature to form -Si-O-H linkages, which can further condense to form -
- 25 Si-O-Si- linkages. The R2 group is preferably a C1-C10 group, and more preferably, a C1-C6 group. The R2 group is preferably an alkyl group such as methyl or ethyl or an alkyl ether group such as 2-methoxyethyl or 2ethoxyethyl
- 30 The linking group L should contain at least one alkoxy group, and preferably a propoxy group, and the silane adhesion promoter is preferably propoxylated. Preferably, the linking group L contains at least one branched propoxy group of the type
- 35 -[OCH2CH(CH2)]- or -[OCH(CH2)CH2]- rather than a linear propoxy group of the type - [OCH2CH2CH2]. Combinations of

propoxy groups of the type
-[OCH₂CH(CH₃)]- or -[OCH(CH₃)CH₂]- can be used.

Preferably, a propoxy group is directly bonded to the radiation-curable group which, preferably, is acrylate.

In first and second preferred embodiments, the propoxy group-containing adhesion promoter is represented by the following formulas:

 $\label{eq:ch2-coo} \mbox{ $\{\mbox{CH}_2\mbox{CH}c\mbox{H}_2\mbox{CH}_2\mbox$

10

 $CH_2=CH-COO\{CH_2CHCH_3O\}_x-COCH_2CH_2Si(OCH_3)_3$ (2)

wherein R³ is H or methyl and X can be about 1-8.

Preferably, X is higher than 1.5, preferably higher

15 than 2. In a currently preferred embodiment, X is equal
to about 4. X may represent a statistical average
wherein different silane molecules have different
levels of propoxylation.

A preferred example of the silane compound 20 is PRO 1779 available from Sartomer.

 $\label{eq:Another example of a suitable compound is} % \[\frac{1}{2} \left(\frac{1}{2} \right) = \frac{1}{2} \left(\frac{1}{2} \right) \left(\frac{1}{$

 $CH_2 = CR^3COO - [CH (CH_3) CH_2O]_x CH_2 CH_2 CH_2 Si (OCH_3)_3$ (3)

25

The amount of the silane adhesion promoter, or mixture thereof, can be about 0.1 wt.* to about 30 wt.*, and preferably, about 0.3 wt.* to about 5 wt.*, and more preferably, about 0.5 wt.* to about 3 wt.*.

30 Small amounts of other types of silane adhesion promoters, which do not include branched propoxy groups, can be included such as ymercaptopropyltrimethoxysilane. However, they are preferably excluded in favor of the propoxylated silane 35 compound. If present, other types of silane adhesion

promoters can be used in amounts of 0.3 wt.% or less, and preferably, 0.1 wt.% or less.

Silane adhesion promoters, or coupling agents, are discussed in E.P. Plueddemann's <u>Silane</u>

<u>Coupling Agents</u>, Plenum Press, New York 1982, which can guide one skilled in the art on the preparation and use of silanes.

Additives are commonly used in radiationcurable coatings, including optical fiber coatings, and
10 known additives can be used in effective amounts. For
example, slip agents may be used to reduce the
coefficient of friction and thermal antioxidants may be
used to improve thermal and oxidative stability. Other
representative additives include stabilizers well-known
15 in the art to prevent gelation, UV screening compounds,
leveling agents, polymerization inhibitors, light
stabilizers, chain transfer agents, colorants including
pigments and dyes, plasticizers, fillers, wetting
improvers, preservatives, and the like. Other polymers
20 and oligomers can be added to the compositions as
needed.

Moisture content in the coatings is preferably minimized.

Known thermal antioxidants can be used to

further increase thermo-oxidative stability. In a

preferred embodiment, the thermal antioxidant such as

Irganox 245 or Irganox 1035 (Ciba Geigy) is used. The

amount of antioxidant can be about 0.1 wt.* to about 5

wt.*, and preferably, about 0.3 wt.* and about 2 wt.*.

For an inner primary coating, cure speed is preferably about 0.5 J/cm^2 or less (at 95% of maximum attainable modulus), and more preferably, about 0.3 J/cm^2 or less, and even more preferably, about 0.2 J/cm^2 or less.

For an outer primary coating or matrix

material, cure speed is preferably about 0.5 J/cm2 or less (at 95% of maximum attainable modulus), and more preferably, about 0.3 J/cm2 or less, and even more preferably, about 0.2 J/cm2 or less.

- For a cured coating composition, the dry adhesion values as determined by the test procedures section described herein range from about 5 to 1000 g.force and the wet adhesion values range from about 1 to 900 g.force. Preferably, both the dry and wet 10 adhesion values are about 5 g.fore or higher and more preferably about 10 g.force or higher. It is also preferred that the dry and wet adhesion values are about 300 g.force or lower, more in particular, 100 g.force or lower.
- 15 For an inner primary coating, the segment modulus is preferably about 0.1 MPa to about 10 MPa. Tensile strength is preferably at least about 0.2 MPa. more preferably, at least about 0.3 MPa and most preferably about .7 to about 3.5. Elongation is
- 20 preferably about 30% or more, and more preferably, about 40% or more. There are no particular upper limits on tensile strength and elongation, but in general. tensile strength will be less than about 10 MPa, and elongation will be less than about 400%.
- 25 For an inner primary coating, refractive index is preferably about 1.47 or above, and more preferably, about 1.48 or above.

Conventional methods in the radiation-cure and optical fiber arts can be used to cure the 30 compositions including electron-beam and UV cure. Thermal cure is less preferred, although some thermal cure may possibly occur under a hot UV lamp. UV cure is preferred. Electron beam cure provides the advantage that photoinitiators may be omitted.

The production of and useful 35 characteristics for coated optical fibers are discussed

20

in, for example, U.S. Patent No. 5,104,433, which is hereby incorporated by reference. Single mode or multimode fiber can be prepared by known methods. Step index and graded index fibers can be prepared by known 5 methods. In the coated fiber, loss due to absorption, scattering, macrobending and microbending should be minimized. Avoiding microbending loss is particularly important. Glass fiber typically is about 125 microns in diameter, and coating layers of approximately 30 microns are applied thereto.

Optical fiber ribbons are described in. for example, U.S. Patent No. 4,900,126 to Jackson et al.: 5,373,578 to Parker et al., 5,379,363 to Bonicel et al.; the complete disclosure of which is hereby incorporated by reference. Ribbon stripping is discussed in, for example:

"Testing of 4- and 8-Fiber Ribbon Strippability" by G.A. Mills in Int. Wire & Cable Symp. Proc., 1992, pqs. 472-474; and

"The Effect of Fiber Ribbon Component Materials on Mechanical and Environmental Performance" by K.W. Jackson et al., in Int. Wire & Cable Symp. Proc., 1993, pgs. 28-34; which are hereby incorporated by reference. Preferably, the compositions are adapted 25 to provide for good ribbon stripping behavior.

In addition to ribbon packaging, fiber designs can include tight buffer, loose tube, filled loose tube, and mini-bundle. Cables can be packaged by conventional buffering, stranding, and jacketing steps.

Optical fiber fabrication is disclosed in, for example, the article "Fiber Optics" Encyclopedia of Chemical Technology, Vol. 10, 4th Ed. 1993, pg. 514-538, which is hereby incorporated by reference.

The invention will be further illustrated 35 by means of the following non-limiting examples.

EXAMPLES

Unless indicated otherwise, radiation-cure was effected in these examples with use of 75 μm films cured under a Fusion D lamp, nitrogen atmosphere, 8 cubic feet per minute flow, at 1.0 J/cm².

A radiation-curable formulation was prepared from pre-mixture ingredients according to Table I. In Table I, the composition of Example 1 comprises a radiation-curable silane adhesion promoter also containing propoxy groups. In contrast, comparative example A comprises the silane adhesion promoter, γ-mercaptopropyltrimethoxy-silane, which does

not contain propoxy groups.

15

TABLE I

-		
COMPONENT (wt.%)	1	A
urethane acrylate oligomer ¹	70.85	70.85
vinyl caprolactam	4	4
phenoxyethyl acrylate	18	18.65
IRGACURE 184 (Ciba)	4.0	4.0
IRGANOX 1076 (Ciba)	0.5	0.5
CYGAURD (CYTEC) AO 711	1.0	1.0
Pro 1779 ²	1.65	
γ-mercaptopropyltrimethoxysilane		1.0
PROPERTIES		
viscosity (mPa.s)	5,800	5,500
tensile strength (MPa)	0.6	0.4
elongation (%)	67	71
secant modulus (MPa)	1.4	1.0
cure speed	0.16	0.5
(J/cm ² @95%cure)		
dry adhesion (g.force)	67	83
wet adhesion (g.force)	25	56

¹urethane acrylate oligomer of a mixture of an acrylic polyol and polypropyleneglycol

2 [CH2=CH-COO{CH2CHCH3O}x-COCH2CH2]2NCH2CH2CH2Si(OCH3)3

5 Examples 2-3

Radiation-curable compositions were formulated from pre-mixture ingredients according to Table II.

10 Table TT

2 (control)	3
65.4	65.30
20.00	19.50
10.00	10.00
3.00	3.00
0.5	
	0.5
	0.1
1.00	
	1.50
0.10	0.10
0.67	0.23
	20.00 10.00 3.00 0.5 1.00

1014-4 is prepared as a mixture of the oligomer, which is prepared from PTG-L2000 polyether polyol, IPDI (isophorone diisocyanate), and HEA (hydroxyethyl 15 acrylate)oligomer, HEA-(IPDI-PTG-L2000), and a monomer, ethoxylated nonylphenolacrylate. The 1014-4 oligomer for example 2 is a mixture of 50.3 wt.% oligomer and 15.1 wt.% monomer and for example 3 is a mixture of 50.2 wt.% oligomer and 15.1 wt.% monomer.

The adhesion of cured samples on a glass plate at 50% relative humidity and 95% relative humidity were tested using a universal testing instrument, INSTRON Model TTD. The load cell had a ten pound capacity. Glass plates, polished, 20 X 20 cm, (Alletch Associates catalog number 26080) were used. The test material was applied to the glass plates and cured at 1 J/ cm² with a Fusion D Lamp under Nitrogen atmosphere. The thickness of the cured film was about 0 75 microns.

The cured films were held at 50% relative humidity, at about 23°C, for seven days prior to testing.

Test specimens, approximately 1 inch in

15 width and 5 inches long, were cut parallel to the
direction in which the drawdown of the cured film was
prepared. A thin layer of talc was applied to the first
and third strips on each drawdown to reduce blocking
during the adhesion test.

The instrument was calibrated prior to 20 testing. The crosshead speed was set to 10.00 inch/min. For each material, the force required to remove four test specimens from the glass plate was measured and recorded on a strip chart recorder. The value reported 25 is the average of the four measured values. The test specimens remaining on the glass plate were then held at 95% relative humidity, at about 23°C, in an environmental chamber for 1 more day. Prior to removing the plates from the environmental chamber, a layer of 30 slurry (fine powdered polyethylene and water) was applied to the surface of the drawdown to retain the moisture. For each material, the force to remove four test specimens from the glass plate was measured as above.

The tensile strength of cured samples was

30

tested using a universal testing instrument INSTRON
Model 4201 equipped with a personal computer and
software "Series IX Materials Testing System." The
load cells used were 2 and 20 pound capacity. The ASTM
D638M was generally followed, with the following
modifications

A drawdown of each material to be tested was made on a glass plate and cured using a UV processor. The cured film was conditioned at 23 \pm 2°C and 50 \pm 5% relative humidity for a minimum of sixteen hours prior to testing.

A minimum of eight test specimens, having a width of .5 \pm 0.002 inches and a length of 5 inches, were cut from the cured film.

To minimize the effects of minor sample defects, sample specimens were cut parallel to the direction in which the drawdown of the cured film was prepared. If the cured film was tacky to the touch, a small amount of talc was applied to the film surface using a cotton tipped applicator.

The test specimens were then removed from the substrate. Caution was exercised so that the test specimens were not stretched past their elastic limit during the removal from the substrate. If any 25 noticeable change in sample length had taken place during removal from the substrate, the test specimen was discarded.

If the top surface of the film was talc coated to eliminate tackiness, then a small amount of talc was applied to the bottom surface of test specimen after removal from the substrate.

The average film thickness of the test specimens was determined. At least five measurements of film thickness were made in the area to be tested (from 35 top to bottom) and the average value used for

calculations. If any of the measured values of film thickness deviates from the average by more than 10% relative, the test specimen was discarded. All specimens came from the same plate.

The appropriate load cell was determined by using the following equation:

[A X 145] X 0.0015 = C

Where: A = Product's maximum expected tensile strength (MPa); 145 = Conversion Factor from MPa to psi; 0.00015

10 = approximate cross-sectional area (in₂) of test specimens; and C = lbs. The 2 pound load cell was used.

specimens; and C = lbs. The 2 pound load cell was used for materials where C=1.8 lbs. The 20 pound load cell was used for materials where 1.8 (C (18 lbs). If C) 19, a higher capacity load cell was required.

- The crosshead speed was set to 1.00 inch/min, and the crosshead action was set to "return at break". The crosshead was adjusted to 2.00 inches jaw separation. The air pressure for the pneumatic grips was turned on and adjusted as follows: set
- approximately 20 psi(1.5 Kg/cm²) for primary optical fiber coatings and other very soft coatings; set approximately 40 psi(3Kg/cm²) for optical fiber single coats; and set approximately 60 psi (4.5 Kg/cm²) for secondary optical fiber coatings and other hard
- 25 coatings. The appropriate INSTRON computer method was loaded for the coating to be analyzed.

After the INSTRON test instrument had been allowed to warm-up for fifteen minutes, it was calibrated and balanced following the manufacturer's operating procedures.

The temperature near the INSTRON Instrument was measured and the humidity was measured at the location of the humidity gage. This was done just before beginning measurement of the first test specimen.

Specimens were only analyzed if the temperature was within the range 23 ± 1.0°C and the relative humidity was within 50 ± 5%. The temperature was verified as being within this range for each test specimen. The humidity value was verified only at the beginning and the end of testing a set of specimens from one plate.

Each test specimen was tested by suspending it into the space between the upper pneumatic grips such that the test specimen was centered laterally and hanging vertically. Only the upper grip was locked. The lower end of the test specimen was pulled gently so that it has no slack or buckling, and it was centered laterally in the space between the open lower grips.

While holding the specimen in this position, the lower grip was locked.

The sample number was entered and sample dimensions into the data system, following the instructions provided by the software package.

20 The temperature and humidity were measured after the last test specimen from the current drawdown was tested. The calculation of tensile properties was performed automatically by the software package.

The values for tensile strength, &
25 elongation, and (secant or segment) modulus were
checked to determine whether any one of them deviated
from the average enough to be an "outlier." If the
modulus value was an outlier, it was discarded. If
there were less than six data values for the tensile
30 strength, then the entire data set was discarded and
repeated using a new plate.

The viscosity was measured using a PHYSICA MC10
Viscometer. The test samples were examined and if an
<u>Viscosity</u> excessive amount of bubbles was present,
35 steps were taken to remove most of the bubbles. Not all

acceptable.

bubbles need to be removed at this stage, because the act of sample loading introduces some bubbles.

The instrument was set up for the

conventional Z3 system, which was used. The samples

were loaded into a disposable aluminum cup by using the
syringe to measure out 17 cc. The sample in the cup was
examined and if it contains an excessive amount of
bubbles, they were removed by a direct means such as
centrifugation, or enough time was allowed to elapse to

let the bubbles escape from the bulk of the liquid.
Bubbles at the top surface of the liquid are

The bob was gently lowered into the liquid in the measuring cup, and the cup and bob were

15 installed in the instrument. The sample temperature was allowed to equilibrate with the temperature of the circulating liquid by waiting five minutes. Then, the rotational speed was set to a desired value which will produce the desired shear rate. The desired value of the shear rate is easily determined by one of ordinary skill in the art from an expected viscosity range of

the sample.

The instrument panel read out a viscosity value, and if the viscosity value varied only slightly (less than 2% relative variation) for 15 seconds, the measurement was complete. If not, it is possible that the temperature had not yet reached an equilibrium value, or that the material was changing due to shearing. If the latter case, further testing at different shear rates will be needed to define the sample's viscous properties. The results reported are the average viscosity values of three test samples.

Cure speed was determined by Dose vs.

Modulus curve analysis of radiation-cured sample films

of approximately 75 microns thickness which were cured
at each of a series of doses (0.2, 0.3, 0.5, 0.75,

1.0, and 2.0). Six specimens were cut from the center portion of each plate of the samples prepared. A Universal Testing Instrument, INSTRON Model 4201 equipped with a suitable personal computer and software "Series IX Materials Testing System" was used to measure the modulus of each specimen. The modulus measurements were then entered into the software package and the calculations were automatically performed with a determination of the average modulus for each film sample. A least squares fit of the modulus versus dose data was then performed to fit the non-linear equation:

modulus = $K_1 \times [1 - e]^{(k2 \times dose)}$

15

This is best done using computer software such as SIGMA PLOT to generate the appropriate constraints.

The dose-modulus curve was then created by plotting the modulus values as a scatter plot and the 20 above equation as a line.

The cure speed of the coating composition is defined as the dose at which 95% of the ultimate secant modulus is attained.

Small sections of the cured coating are
placed on a microscope slide under a cover slip. These
sections are immersed in one of a series of liquids of
known refractive index and the resulting preparation
observed through a microscope. The characteristic
optical phenomenon known as the Becke' line is used to
determine the extent and direction of the mismatch
between the liquid and solid. Additional trials are
made with other liquids from the series of known
refractive indices until a match is found and the
refractive index value determined.

While the invention has been described in detail and with reference to specific embodiments

thereof, it will be apparent to those of ordinary skill in the art that various changes and modifications can be made therein without departing from the spirit and scope thereof.

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WHAT IS CLAIMED IS:

- A radiation-curable composition for use as an optical fiber material or coating comprising the combination of the following pre-mixture incredients:
 - (A) about 5 wt.% to about 95 wt.% of one or more radiation-curable oligomers,
 - (B) about 5 wt.% to about 95 wt.% of one or more monomer diluents,
 - (C) optionally, one or more photoinitiators,
 - (D) about 0.1 wt.* to about 30 wt.* of one or more adhesion promoters represented by: $R^1\!-\!L\!-\!Si\left(OR^2\right)_3$
- wherein R^1 is a group comprising an ethylenically unsaturated radiation-curable group; said three R^2 groups independently of each other are C_1 - C_{10} groups; and L is a linking group which comprises one or more branched alkoxy groups.
- A radiation-curable composition according to claim 1, wherein said group L comprises two or more branched alkoxy groups.
- A radiation-curable composition according to any one of claims 1-2, wherein said group L comprises
 at least one branched propoxy group.
 - A radiation-curable composition according to any one of claims 1-3, wherein said group L further comprises an amino propyl groups.
- A radiation-curable composition according to any
 one of claims 1-4, wherein said R group comprises an acrylate or methacrylate group.
 - A radiation-curable composition according to any one of claims 1-5, wherein R² is a methyl or ethyl group.
- A radiation-curable composition according to any one of claims 1-6, wherein said composition is

formulated to be an inner primary coating having suitable adhesion to glass and a cure speed at 95% cure of at least 0.5 J/cm² or less.

- A radiation-curable composition according to claim 7, wherein said cure speed is at least 0.3 J/cm² or less measured from dose-modulus curves.
 - A radiation-curable composition according to any one of claims 1-8, wherein the dry and wet adhesion are higher than 5, and lower than 300
- 10 g.force.

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- A radiation-curable composition according to any one of claims 1-9, wherein said composition is cured to attain at least 80% of its maximum attainable modulus.
- 15 11. A coated optical fiber comprising an optical fiber for transmitting light and at least one composition according to any one of claims 1-10.
 - 12. The coated optical fiber according to claim 11, wherein the coating has a tensile strength of 0.2
- 20 MPa or higher and about 10 MPa or lower.

 13. Coated optical fiber according to any one of
 - Coated optical fiber according to any one of claims 11-12, wherein the coating has an elongation at break of 30-400%.
 - 14. A method for coating optical fiber which
- comprises the steps of: drawing an optical fiber; applying one or more radiation-curable inner primary, outer primary, or single coating compositions; said coating composition in a precured state being a composition according to any
 - one of claims 1-10 and suitably curing said one or more radiationcurable coating compositions.

INTERNATIONAL SEARCH REPORT

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A. CLASSIFICATION OF SUBJECT MATTER
TPC 6 C03C25/02 C08K5/54 C08K9/06 G02B1/04 C09D201/10 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED mum documentation searched. (classification system followed by classification symbols) Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Category * Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. US 4 849 462 A (BISHOP TIMOTHY E) 1-14 Α 18 July 1989 cited in the application see the whole document US 5 336 563 A (COADY CLIVE J ET AL) 1 - 14A 9 August 1994 cited in the application see the whole document US 4 472 019 A (BISHOP TIMOTHY E ET AL) 1-14 Α 18 September 1984 cited in the application see the whole document WO 95 06650 A (DSM NV) 9 March 1995 1-6 Α cited in the application see the whole document -/--Further documents are listed in the continuation of box C. Patent family members are listed in annex. "T" later document published after the international filing date or priority date and not in conflict with the application but clied to understand the principle or theory underlying the revention. Special categories of cited documents : "A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier document but published on or after the International filling date "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document le taken alone "L" document which may throw doubts on priority claim(a) or which is cited to establish the publication date of another citation or other special reason (as specified) **NOTE SIL INVESTIVE SEEP WHEN THE GOCUMENT IE LIKEN BIODE.

"O document of particular reviewnce; the claimed Investion
cannot be considered to involve en inventive step when the
document is combined with one or more other such documents, such combination being obvious to a person skilled
in the art. "O" document referring to an oral disclosure, use, exhibition or other means document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the ectual completion of the international search Date of mailing of the international search report 1 February 1999 09/02/1999

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Somann, K

INTERNATIONAL SEARCH REPORT

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